Liver-targeted liposomes for codelivery of curcumin and combretastatin A4 phosphate: preparation, characterization, and antitumor effects

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Background: Recent efforts have been focused on combining two or more therapeutic approaches with different mechanisms to enhance antitumor therapy. Moreover, nanosize drug-delivery systems for codelivering two drugs with proapoptotic and antiangiogenic activities have exhibited great potential in efficient treatment of cancers.

Methods: Glycyrrhetinic acid (GA)–modified liposomes (GA LPs) for liver-targeted codelivery of curcumin (Cur) and combretastatin A4 phosphate (CA4P) were prepared and characterized. In vitro cellular uptake, cytotoxicity, cell migration, in vivo biodistribution, antitumor activity, and histopathological studies were performed.

Results: Compared with unmodified LPs (Cur-CA4P LPs), Cur-CA4P/GA LPs were taken up effectively by human hepatocellular carcinoma cells (BEL-7402) and showed higher cytotoxicity than free drugs. In vivo real-time near-infrared fluorescence–imaging results indicated that GA-targeted LPs increased accumulation in the tumor region. Moreover, Cur-CA4P/GA LPs showed stronger inhibition of tumor proliferation than Cur, Cur + CA4P, and Cur-CA4P LPs in vivo antitumor studies, which was also verified by H&E staining.

Conclusion: GA-modified LPs can serve as a promising nanocarrier for liver-targeted co-delivery of antitumor drugs against hepatocellular carcinoma.

Keywords: liver-targeted, LPs, curcumin, combretastatin A4 phosphate, combination therapy

Introduction
Hepatocellular carcinoma (HCC) is one of the most common malignant tumors, with high incidence and mortality worldwide.1 Chemotherapy is one of the main strategies in the clinical treatment of HCC. However, traditional cytotoxic chemotherapeutic drugs, such as doxorubicin and paclitaxel, exhibit limited clinical efficacy, due to systemic toxicity, lack of selectivity, and drug resistance.2,3 The effectiveness of combination therapy is supported by clinical research, which shows superior antitumor efficacy than single-drug therapy via different antitumor approaches.4,5 Antiangiogenesis, a promising antitumor strategy blocking the development of tumor blood vessels, is widely accepted.6 Therefore, the combination of chemotherapeutics with antiangiogenic drugs is expected to improve therapeutic efficacy for HCC.7,8,9

Curcumin (Cur), a diphenolic compound derived from the rhizome of turmeric, exhibits potent antitumor efficacy against prostate, breast, and colon cancers and belongs to the third generation of anticancer drugs.10 Cur blocks NFκB signaling to inhibit proliferation, thereby inhibiting cancer-cell growth and inducing apoptosis.11,12 In the clinic, single-drug therapy may increase the occurrence of drug resistance, suggesting the necessity of using other active drugs in combination with Cur to
overcome this defect. Hydrophilic combretastatin A4 phosphate (CA4P), isolated from *Combretum caffrum*, is a new antitumor compound targeting blood vessels and soluble phosphate produg derivatives of CA4. CA4P acts mainly on the vascular endothelial cells of the tumor, which causes tumor vasoconstriction, dysfunction, and death of neoplastic cells, due to insufficient blood supply. Few reports about the combination of Cur and CA4P exist. Based on the reduced incidence of drug resistance and tumor treatment in different pharmacological pathways by promoting apoptosis and antitumor angiogenesis, we selected the combination of Cur and CA4P to verify the enhanced antitumor efficacy. The combination of Cur and CA4P can improve the effect of chemotherapy via different mechanisms of action and reduce side effects. However, chronic single administration is often associated with undesirable side effects or drug resistance, which may cause failure in cancer therapy. Possible reasons involve the difficulty in targeted codelivery of the two drugs to tumors, due to their diverse physicochemical properties and mechanisms.

Drug-delivery systems provide a promising opportunity to overcome the dilemma in effective codelivery of multiple cargoes. Many drug-delivery systems, such as liposomes (LPs), microspheres, nanoemulsions, polymer micelles, polymer–drug conjugates, carbon nanotubes, mesoporous silica nanoparticles, have been widely investigated. Among the different nanocarriers, LPs have attracted increased attention. LPs contain a hydrophilic core and hydrophobic shell, which can load hydrophilic, hydrophobic, and amphipathic drugs and improve their solubility. In addition, the flexibility of the membranal structure in LPs would be beneficial for sustained or controlled drug release. Moreover, LPs <200 nm can improve accumulation in solid tumors, due to the enhanced permeability and retention (EPR) effect, which contributes to enhanced antitumor efficacy and reduced systemic toxicity. On the basis of these properties, we selected LPs as nanocarriers to improve therapeutic effects against tumors.

Active liver-targeting LPs must improve the accuracy of drug actions on target sites. Glycyrrhetinic acid (GA) is the active aglycone of glycyrrhizin, which belongs to a pentacyclic triterpene derivate. Specific receptors for GA on the cytomembrane of hepatocytes have been verified. Protein kinase Cα is the target binding protein of GA that shows higher expression in HCC cells than that in adjacent nontumor liver cells. As such, GA-modified LPs can be expected to exhibit increased active targeting to HCC cells and tissue, which will be beneficial for further enhanced antitumor efficacy than unmodified LPs.

In summary, novel GA-modified LPs for effective codelivery of Cur and CA4P (Cur-CA4P/GA LPs) were developed in this study. Cur-CA4P/GA LPs targeted tumor sites effectively and inhibited tumor-cell proliferation and destroyed tumor neovascularization simultaneously, consequentely improving antitumor activity for HCC. In addition to characterization of physicochemical properties of the codelivered LPs, evaluation of in vitro cytotoxicity, cellular uptake, cell migration, and in vivo biodistribution and antitumor efficacy was conducted in this study.

**Methods**

**Materials**

Cur (molecular weight [MW] 368.38 Da), L-α-phosphatidylcholine, cholesterol (Chol; MW 386.67 Da), Sephadex TMG50 medium (separation range 1,000–30,000), MTT, DAPI, RPMI-1640 medium, and FBS were purchased from Solarbio (Beijing, China). GA (MW 470.69 Da) was acquired from Fujie Pharmaceutical (Sanyuan, China). 1,4-(4,6-Dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride (DMTMM) was purchased from Medpep (Shanghai, China). 1-Ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) was purchased from Aladdin Reagent (Shanghai, China). DiR (1,1′-dioctadecyl-3,3,3,3-tetramethyl indotricarbocyanine Iodide) was obtained from Pan Bo Biochemistry (Beijing, China). 1,2-Distearyl-sn-glycerol-3-phosphoethanolamine (DSPE)–N-hydroxysuccinimide (NHS)–polyethylene glycol 2,000 (PEG2,000) and DSPE-PEG2,000–fluorescein isothiocyanate (FITC) were purchased from Ruixi Technology (Xi’an, China), and CA4P (MW 440.30 kDa) was obtained from Innochem Technology (Beijing, China). All chemical reagents used in the experiments were of analytical reagent grade.

**Cell lines**

Human HCC cells (BEL7402) and mouse melanoma cells (B16) were purchased from Wuhan University Life Collection Center (Wuhan, China). BEL7402 cells were cultured in RPMI 1640 medium supplemented with 10% FBS and 100 IU/mL penicillin–streptomycin at 37°C in a humidified incubator containing 5% CO2. B16 cells were cultured with DME:F12 (1:1) medium with 10% FBS and 100 IU/mL penicillin–streptomycin in the same environment. Cells were subcultured with trypsin for ~2 minutes, and the culture medium containing FBS was then added to terminate the digestion of trypsin. Suspensions were centrifuged at 1,000 rpm for 5 minutes and the supernatant discarded. Cells were then counted and inoculated. Murine HCC cells (H22)
were obtained from KeyGen Biotech (Nanjing, China). H22 cells were cultured under the same conditions as BEL7402 cells and used to establish the tumor-bearing mouse model.

Animals
Male BALB/c mice (20±2 g) aged 4–6 weeks were purchased from Pengyue Experimental Animal Breeding (Jinan, China). Animals were fed a standard diet and allowed water ad libitum. All experiments were performed in compliance with the animal management rules of the Ministry of Health of China (document 55, 2001) and the animal experiment ethics review committee of Weifang Medical University (2017-025).

Synthesis of DSPE-PEG$_{2,000}$-GA
DSPE-PEG$_{2,000}$-GA was synthesized by a two-step reaction.$^{31,32}$ In the first step, GA solution in methanol was activated by DMTMM to form the intermediate product – GA-ES. At room temperature, GA was added slowly to an ethylene diamine solution to obtain diamine-modified GA (GA-N). In the second step, GA-N and DSPE-PEG$_{2,000}$-NHS at a molar ratio of 3:1 was dissolved in dimethyl sulfoxide in the presence of EDC (1.58 mg) and 20 µL triethylamine was added to the reaction system to regulate pH. The whole system was reacted for 48 hours under nitrogen protection to prevent bubble formation, followed by dialysis in 1,000 mL double-distilled water for 3 days. The solution was freeze-dried to obtain DSPE-PEG$_{2,000}$-GA. Chemical structures of DSPE-PEG$_{2,000}$-NHS, GA-N, and DSPE-PEG$_{2,000}$-GA were confirmed by $^1$H nuclear magnetic resonance (JNM ECP-600; Jeol, Tokyo, Japan) by dissolving the target molecule in deuteron chloroform.

Preparation of liver-targeted LPs for coloading Cur and CA4P
Cur-CA4P/GA LPs were prepared via the well-established technique of thin-film evaporation.$^{33}$ 1-L-α-Phosphatidylethanol (120 mg) and Chol (40 mg) were mixed in 5 mL chloroform at a mass ratio of 3:1 and 20 mg DSPE-PEG$_{2,000}$-GA was added to this system at a phospholipid:DSPE-PEG$_{2,000}$-GA mass ratio of 6:1 and mixed with 3 mL of 1 mg/mL Cur anhydrous ethanol solution. The mass ratio of Cur and phospholipids was 1:40. Chol was then removed using a rotary evaporator to form a dry-lipid film at 35°C under uniform speed. CA4P dissolved in PBS (pH 7.4, 5 mL) was added to the lipid film, hydrated for 30–60 minutes, and sonicated for 5 minutes to decrease the particle size. The final concentrations of Cur and CA4P in the system were 0.06 and 0.75 mg/mL, respectively. Finally, 5 mL LP solution was obtained. Unencapsulated Cur was removed using Sephadex G50.$^{34}$ The chromatographic column was filled with Sephadex G50 medium. After the chromatographic column had reached equilibrium, the sample was added and eluted with PBS. Dialysis was used to remove the unencapsulated CA4P. The sample was sealed in a dialysis bag, and the bag placed in 500 mL PBS solution for 3 days. Finally, the Cur-CA4P/GA LPs were refrigerated (Figure 1). The whole preparation process was conducted without light.

Single-load CA4P LPs were prepared through reverse-phase evaporation:$^{35}$ 120 mg L-α-phosphatidylcholine and 15 mg Chol were mixed with 8 mL ether at a mass ratio of 8:1; 10 mg CA4P was dissolved in 5 mL PBS, and the former mix added. Both the ether and PBS solutions were mixed and followed by sonication for 1 minute in an ice bath to form a w:o uniform emulsion. Decompression rotating evaporation

![Figure 1 Schematic diagram of Cur-CA4P/GA LPs.](https://www.dovepress.com/)

**Notes:** Cur (green) and CA4P (red) were encapsulated into the hydrophobic shell and hydrophilic core of the LPs, respectively. GA was linked to the LPs by conjugation with DSPE-PEG$_{2,000}$.

**Abbreviations:** CA4P, combretastatin A4 phosphate; Cur, curcumin; LPs, liposomes; GA, glycyrrhetinic acid; DSPE, 1,2-distearoyl-α-glycero-3-phosphoethanolamine; PEG, polyethylene glycol.
removed organic solvents and achieved a colloidal state. Exactly 5 mL PBS was then added for hydration, followed by vortexing to form a uniform suspension. Blank LPs were prepared using the same procedure without drugs. A single-factor method was adopted for prescription screening.

Preparation of FITC-labeled LPs
FITC-labeled LPs were prepared through thin-film evaporation similarly to Cur-CA4P/GA LPs. To begin, 120 mg 1-α-phosphatidylcholine, 40 mg Chol, and 12 mg DSPE-PEG<sub>2,000</sub>-FITC were mixed in 5 mL chloroform at a mass ratio of 30:10:3. Chol was evaporated using a rotary evaporator to form a dry-lipid film at 35°C under 18 rpm. Subsequently, 5 mL PBS was added for hydration, lasting for 30–60 minutes. The final step was sonication for 5 minutes to obtain FITC LPs. The preparation of FITC-GA LPs was similar to that of FITC LPs. The only difference was that 20 mg DSPE-PEG<sub>2,000</sub>-GA was added to the raw materials (mass ratio of 1-α-phosphatidylcholine and DSPE-PEG<sub>2,000</sub>-GA 6:1). The final concentration of FITC LPs was 2.4 mg/mL.

Preparation of DiR-loaded GA LPs
The preparation of DiR-loaded GA LPs was similar to the previous method. First, 120 mg 1-α-phosphatidylcholine, 40 mg Chol, 20 mg DSPE-PEG<sub>2,000</sub>-GA, and 0.2 mg DiR were mixed in 5 mL Chol at a mass ratio of 600:200:100:1 in an eggplant-shaped flask, dried until a thin-lipid film had formed on the rotary evaporator under reduced pressure, and heated in a 35°C water-bath. The film was hydrated with 5 mL PBS, followed by sonication. The obtained DiR-GA LPs were filtered in a 220 nm filter device, and the final concentration of DiR was 40 μg/mL.

Characterization of Cur-CA4P/GA LPs
The morphology of the Cur-CA4P/GA LPs was examined using a transmission electron microscopy. Samples (30 μL) were prepared by placing diluted LPs on a copper grid for 3 minutes, followed by using filter paper to absorb excess samples and staining with 2% phosphotungstic acid (pH 6.5) for 2 minutes. Size distribution and ζ-potential of the LPs were determined by dynamic light scattering at 25°C with a Malvern Zetasizer Nano ZS90. Samples were diluted in PBS and measured in triplicate. For measurement of loading content (LC) and encapsulation efficiency (EE), 1 mL Cur-CA4P/GA LPs was sampled after the removal of free drug and the volume made up with methanol to 10 mL. The absorbance of Cur and CA4P was analyzed with ultraviolet-visible spectrophotometry at 425 and 295 nm. LC and EE were calculated:

\[
LC = \frac{W_{\text{loaded drug}}}{W_{\text{a}}} \times 100\%
\]

\[
EE = \frac{W_{\text{loaded drug}}}{W_{\text{total drug}}} \times 100\%
\]

where \(W_{\text{loaded drug}}\) is drug loaded in LPs, \(W_{\text{total drug}}\) feeding weight of drug, and \(W_{\text{a}}\) weight of all materials.

Stability studies
Cur-CA4P/GA LPs were placed in a 4°C refrigerator for 1, 3, 7, 9, 11, and 14 days. Samples were diluted in PBS to 1 mg/mL for measurement of particle size and ζ-potential, and LC and EE were determined by the sample solution.

In vitro Cur and CA4P release from LPs
In vitro release of Cur and CA4P from LPs was studied by dialysis using PBS (pH 7.4) containing 1% Tween 80 and 20% anhydrous ethanol as release medium. Cur-CA4P/GA LPs (1 mL) suspension was transferred into the dialysis bag (cutoff 3.5 kDa), and the bag was placed into a conical bottle containing 50 mL release solution. The system was kept in a shaking incubator (THZ-82; Jintan Medical instrument Factory, Jintan, China) with stirring speed of 100 rpm at 37°C. After 0.5, 1, 2, 4, 8, 12, 24, 36, and 48 hours, 4 mL medium was removed and the same volume of release medium added. Concentrations of Cur and CA4P were measured with ultraviolet-visible spectrophotometry at wavelengths of 425 and 295 nm, respectively, and their cumulative release was calculated. Cumulative Cur- and CA4P-release percentage were calculated as follows:

\[
Er(\%) = \frac{V \sum_{i=1}^{n-1} C_i + V_0 C_n}{M_d} \times 100\%
\]

where \(M_d\) is the amount of drugs in the LPs, \(V_0\) the whole volume of the release solution, \(C\) the concentration of Cur or CA4P in the release solution, and \(V_r\) the volume of the replaced solution.

In vitro cellular uptake assays
Cellular uptake of liposomal formulations was evaluated in BEL7402 cell by qualitative methods. Cells (5×10<sup>4</sup>) were seeded onto a glass-bottom cell-culture dish (φ20 mm) and
incubated for 24 hours at 37°C until adherent growth was observed. After incubation with FITC LPs or FITC-GA LPs (2.4 mg/mL) for 30 minutes, all reagents were removed and cells washed in cold PBS (pH 7.4) three times, fixed with 4% polyformaldehyde for 10 minutes, and stained with DAPI (0.1 µg/mL, diluted with PBS) for 5 minutes in the CO₂ incubator. Finally, cells were washed three times with PBS and photographed under confocal laser-scanning microscopy (TCSSP8; Leica, Wetzlar, Germany). Excitation and emission wavelengths of DAPI and FITC were 359 and 461 nm and 490 and 520 nm, respectively. All operations were performed in the dark. Images were merged with Image-Pro Plus 6.0.

**In vitro cytotoxicity assays**

The in vitro cytotoxicity of different formulations was assessed using MTT assays. Approximately 5x10⁵ B16 or BEL7402 cells were seeded in 96-well plates and incubated for 24 hours at 37°C. Cell-compatibility tests had to be performed before cells were treated with drugs. Cells were treated with different concentrations of blank GA LPs for 24 or 48 hours at 37°C. MTT reagent (10 µL, 5 mg/mL) was added to each well and incubated for 4 hours at 37°C in the dark. Approximately 150 µL dimethyl sulfoxide was then added to each well and shaken for 10 minutes to dissolve the formazan crystals. The absorbance value of each hole was detected at 490 nm using an enzyme marker (ELX800; BioTek Instruments, Winooski, VT, USA). Cells were harvested. The in vitro cytotoxicity of different formulations was calculated using MTT assays.

**Cell-migration assays**

Cell migration of free Cur and CA4P, Cur + CA4P, Cur-CA4P LPs, and Cur-CA4P/GA LPs were assessed through wound-healing assays. In brief, 10⁴ B16 or BEL7402 cells were seeded in six-well plates and grown for 24 hours at 37°C in a CO₂ incubator. When cell density reached 90%, cells were scratched with a sterile pipette tip with at least three lines per well. Scratched cells were washed off with PBS and the different drug formulations added. Images of cells invading the scratch were captured at 0, 24, and 48 hours using microscopy. The effect of drugs on cell migration was evaluated by calculating the migration rate:

\[
\text{Migration rate} = \left(1 - \frac{W_n - W_0}{W_0}\right) \times 100\%
\]

where Wₙ and W₀ represent average scratch widths at n and 0 hours, respectively.

**In vivo real-time near-infrared fluorescence imaging**

Near-infrared fluorescence imaging (NIRF) was used to observe the biodistribution of GA LPs formulation in vivo, and DiR-GA LPs (40 µg/mL) were prepared to monitor the fate of LPs. When tumors were 100 mm³, DiR-GA LPs were injected into tail veins of mice. Free DiR was used as control. Precisely 10 mg DiR iodide in a 10 mL volumetric bottle was dissolved in methanol at a constant volume and prepared into a 1 mg/mL drug-reserve solution, which was stored at 4°C in the refrigerator for later use. DiR iodide solution was diluted with normal saline to the concentration of 40 µg/mL. Mice were anesthetized with 10% chloral hydrate and real-time NIRF images obtained with excitation and emission at 745 and 835 nm, respectively. Results were analyzed using the Living Image 3.1 software (PerkinElmer, Waltham, MA, USA).

**In vivo antitumor efficacy**

To evaluate the in vivo antitumor efficacy of the Cur-CA4P/GA LPs, we established the H22 tumor-bearing BALB/c male mouse model. Establishment of this model was divided into two steps. First, 0.3 mL hepatoma H22 cells was injected into the abdominal cavity of five mice. After 2 weeks, the mice had developed ascites tumors. Second, cells from the H22 liver cancer ascites-tumor sources were collected in sterile conditions and diluted to 4x10⁶ tumor cells/mL with normal saline. Approximately 0.2 mL of the suspension was subcutaneously inoculated into the flanks of mice. When tumor volume had reached ~100 mm³, mice were divided into five groups: normal saline (control group), free Cur, Cur + CA4P, Cur-CA4P LPs, and Cur-CA4P/GA LPs. Mice in each preparation group were administered 5 mg/kg Cur and 25 mg/kg CA4P every 2 days. During the treatment course, mice were weighed and diameters of tumors measured. Tumor volumes were calculated according to the formula \( V = \frac{4}{3} \pi R^2 \times L \) in which \( L \) is the longest and \( W \) the shortest tumor diameter (mm). Mice were killed and tumors harvested after completion of 2 weeks of treatment. Tumor weights were measured, and tumor growth–inhibition rate was calculated: \( (1 - \text{tumor weight of drug-treated group/tumor weight of control group}) \times 100\% \).

**Histochemical staining**

After the drug administration in the in vivo antitumor study, tumor tissue samples were removed, fixed in 4% formaldehyde for >24 hours, and embedded in paraffin. Paraffin-embedded tumor samples were cut into
slices of 4 µm using an ultrathin slicer (KD3358; Cody Instrument Equipment). For histological evaluation, H&E was used for tumor-tissue staining. Images were magnified 400 times by observation and photography with positive fluorescence microscopy (IX51; Olympus, Tokyo, Japan).

Statistical analyses
All data are expressed as means ± SD. Student’s t-test was used to determine whether differences between test groups were statistically significant. P<0.05 was considered statistically significant.

Results and discussion
Synthesis and characterization of DSPE-PEG$_{2,000}$-GA
The targeting material DSPE-PEG$_{2,000}$-GA was synthesized by coupling aminated GA to the DSPE-PEG$_{2,000}$-NHS. Figure 2 indicates that GA was first modified by ethylenediamine to form GA-N. Second, the free-amino group in GA-N was reacted with DSPE-PEG$_{2,000}$-NHS to generate DSPE-PEG$_{2,000}$-GA. Figure 3 shows the characteristic peaks of DSPE-PEG$_{2,000}$-NHS at 3.6 ppm (PEG group) in the NMR spectra obtained. Peaks at 0.7–1.5 ppm were assigned to the triterpenoid protons of GA-N, and peaks of

![Diagram of GA-N and DSPE-PEG$_{2,000}$-GA synthesis](image)

**Figure 2** Synthesis of GA-N (A) and DSPE-PEG$_{2,000}$-GA (B).

**Abbreviations:** GA-N, diamine-modified glycyrrehetic acid; DSPE, 1,2-distearoyl-sn-glycero-3-phosphoethanolamine; PEG, polyethylene glycol; NHS, N-hydroxysuccinimide; EDC, 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide.
Figure 3 1H NMR spectra of DSPE-PEG<sub>2,000</sub>-NHS, GA-N, and DSPE-PEG<sub>2,000</sub>-GA.

Note: Peaks of GA-N at 0.7–1.5 ppm (a) and peaks of DSPE-PEG<sub>2,000</sub>-NHS at 3.6 ppm (b).

Abbreviations: NMR, nuclear magnetic resonance; DSPE, 1,2-distearoyl-sn-glycero-3-phosphoethanolamine; PEG, polyethylene glycol; NHS, N-hydroxysuccinimide; GA-N, diamine-modified glycyrrhetinic acid.

DSPE-PEG<sub>2,000</sub>-GA were observed at 3.6 and 0.7–1.3 ppm. These results showed that GA-N had been successfully introduced into DSPE-PEG<sub>2,000</sub>-NHS by the presence of characteristic peaks at 3.6 and 0.7–1.3 ppm in the DSPE-PEG<sub>2,000</sub>-GA.

Characterization of liver-targeting LPs for codelivery of Cur and CA4P

Mean particle size, ζ-potential, EE, and LC of each LP formulation are shown in Figure 4. LC and EE were measured when the ratios of Cur to CA4P ranged from 1:1 to 10:1, and the third group (Cur:CA4P 1:5) was selected as the representative candidate for further research, owing to its higher LC and EE values compared to those of other groups. Cur-CA4P/GA LPs were prepared by membrane dispersion (Figure 1). LPs were composed of a hydrophilic core and hydrophobic shell. The hydrophobic Cur was loaded into the hydrophobic membranes, whereas CA4P was encapsulated in the hydrophilic core. DSPE-PEG<sub>2,000</sub>-GA was embedded in the outer membrane. Morphological characteristics of the LPs were observed by transmission electron microscopy. Cur-CA4P/GA LPs showed a circular bubble structure with uniform size, which was similar to that of the blank LPs (Figure 5A and B). As shown in Figure 5C, Cur-CA4P/GA LPs showed an average particle size of ~166 nm with narrow size distribution (polydispersity index 0.23). LPs <200 nm in size are effectively easily accumulated in tumor via the EPR effect. The surface charge of LPs is an indication of their stability and interaction with cells. Table 1 shows that...
Figure 4 Characterization of LPs at various Cur:CA4P ratios.
Notes: Particle size, ζ-potential (A), encapsulation efficiency (B), and loading content (C) of the LPs. Values expressed as means ± SD.
Abbreviations: LPs, liposomes; Cur, curcumin; CA4P, combretastatin A4 phosphate; DLS, dynamic light scattering.

Figure 5 Characteristics of blank LPs and Cur-CA4P/GA LPs.
Notes: (A, B) Transmission electron microscopy; (C) particle-size distribution of Cur-CA4P/GA LPs.
Abbreviations: LPs, liposomes; Cur, curcumin; CA4P, combretastatin A4 phosphate; GA, glycyrrhetinic acid.
Table 1 Characterization of various LP formulations

<table>
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<th>DLS (nm)</th>
<th>PD1</th>
<th>ζ-potential (mV)</th>
<th>EE (%)</th>
<th>LC (%)</th>
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<tr>
<td>Blank</td>
<td>97.27±5.51</td>
<td>0.14±0.01</td>
<td>−15.4±2.63</td>
<td>−</td>
<td>−</td>
</tr>
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<td>Cur LPs</td>
<td>135.55±4.83</td>
<td>0.15±0.01</td>
<td>−30.04±2.82</td>
<td>77.4±2.61</td>
<td>1.48±0.18</td>
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<tr>
<td>CA4P LPs</td>
<td>110.92±6.25</td>
<td>0.14±0.04</td>
<td>−28.1±4.52</td>
<td>87.03±1.84</td>
<td>5.48±0.13</td>
</tr>
<tr>
<td>Cur-CA4P LPs</td>
<td>159.45±6.32</td>
<td>0.23±0.01</td>
<td>−26.77±3.85</td>
<td>84.15±1.36 (CA4P)</td>
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<tr>
<td>Cur-CA4P/GA LPs</td>
<td>166.05±4.64</td>
<td>0.23±0.01</td>
<td>−25.67±3.19</td>
<td>84±1.37 (CA4P)</td>
<td>5.25±0.18 (CA4P)</td>
</tr>
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Note: Values expressed as means ± SD.

Abbreviations: LP, liposome; DLS, dynamic light scattering; PDI, polydispersity index; EE, encapsulation efficiency; LC, loading content; Cur, curcumin; CA4P, combretastatin A4 phosphate; GA, glycyrrhetinic acid.

the ζ-potential of Cur-CA4P LPs and Cur-CA4P/GA LPs was −25 mV and exhibited good stability. The stability of Cur-CA4P/GA LPs was investigated for 2 weeks, and no significant changes in particle size, ζ-potential, EE, and LC were observed (P>0.05, Figure 6).

Cur and CA4P release from LPs in vitro

In vitro drug release of Cur and CA4P from Cur-CA4P/GA LPs was evaluated in PBS (pH 7.4) containing 1% Tween 80 and 20% anhydrous ethanol at 37°C. As shown in Figure 7, no obvious burst release of either drug was observed. Both drugs showed sustained release within 48 hours. The results showed certain sustained-release effects of agents encapsulated in the LP system. Cumulative release rates for both drugs were almost the same during the 10 hours of the experiment, whereas CA4P released more quickly than Cur after 10 hours. Approximately 60% of the CA4P was released within 24 hours, whereas <50% of Cur was released during

Figure 6 Characterization of Cur-CA4P GA LP formulations.

Notes: Size distribution and ζ-potential of Cur-CA4P/GA LPs; (A) EE and LC of Cur (B) and CA4P (C) for 2 weeks. Values expressed as means ± SD.

Abbreviations: Cur, curcumin; CA4P, combretastatin A4 phosphate; GA, glycyrrhetinic acid; LPs, liposomes; EE, encapsulation efficiency; LC, loading content; DLS, dynamic light scattering.
Figure 7 In vitro release profiles of Cur and CA4P from Cur-CA4P/GA LPs.

Notes: Release medium: PBS buffer (pH 7.4; 1% Tween 80 and 20% anhydrous ethanol), temperature 37°C. Data expressed as mean ± SD (n=3).

Abbreviations: Cur, curcumin; CA4P, combretastatin A4 phosphate; GA, glycyrrhetinic acid; LPs, liposomes.

Figure 8 Fluorescence microscopy of BEL7402 cells incubated with FITC LPs (A) and FITC-GA LPs (B).

Note: FITC (green fluorescence) and DAPI channels for nucleus (blue fluorescence) were presented.

Abbreviations: FITC, fluorescein isothiocyanate; LPs, liposomes; GA, glycyrrhetinic acid.

In vitro release profiles of Cur and CA4P from Cur-CA4P/GA LPs. The possible explanation is that the GA LPs were taken up by the cancer cells via GA-receptor-mediated endocytosis. The results suggest that the GA molecule may promote the uptake of LPs, resulting in additional fluorescence signal accumulated in the cytoplasm.

In vitro cytotoxicity assays

First, MTT was used to investigate the cytotoxicity of blank GA LPs against BEL7402 and B16 cells. Figure 9A and B show that cellular viability was >86% after blank GA-LP treatment for 24 hours. This result suggested that blank GA LPs exhibited no obvious toxicity at experimental conditions and can be used as drug-delivery carriers.

In vitro cytotoxicity testing of drug-loaded LPs was conducted against BEL7402 and B16 cells. As shown in Figure 9A, free Cur, CA4P, and Cur + CA4P, Cur-CA4P LPs, and Cur-CA4P-GA LPs expressed dose-dependent cytotoxic effects in BEL7402 cells after incubation for 24 hours, which was similar to that in B16 cells (Figure 9B). Similar to the free-drug combination, codelivery of LP formulations exhibited time-dependent cytotoxicity against the two cell lines for 24, 48, and 72 hours (Figure 9C and D). Compared with free Cur and CA4P, the combination of Cur and CA4P showed higher cytotoxicity against BEL7402 and B16 cells, as shown in Figure 9A and B. The result suggested that the combination of chemotherapy drugs with antiangiogenic drugs exhibited cytotoxicity. Moreover, as shown in Figure 9C and D, Cur-CA4P LPs displayed stronger cytotoxicity than the free mixture of Cur and CA4P. Interestingly, Cur-CA4P/GA LPs...
Figure 9 Viability of cells treated with various combinations.

Notes: (A, B) Blank GA LPs, free Cur, CA4P, free Cur + CA4P, Cur-CA4P LPs, or Cur-CA4P/GA LPs in BEL7402 and B16 cells for 24 hours. (C, D) Viability of cells treated with various formulations against BEL7402 and B16 cells for 24, 48, and 72 hours. *P < 0.05, **P < 0.01 (A, B). *Cur-CA4P LPs vs free Cur + CA4P; **Cur-CA4P/GA LPs vs free Cur + CA4P (C, D).

Abbreviations: GA, glycyrrhetinic acid; LPs, liposomes; Cur, curcumin; CA4P, combretastatin A4 phosphate.

showed higher cytotoxicity compared with Cur-CA4P LPs (P < 0.05). This was because the introduction of GA increased cellular uptake of drug-loaded LPs, leading to additional accumulation of drugs in tumor cells.

Cell-migration assays
To confirm the cytotoxic effect of various formulations further, we performed wound-healing assays in BEL7402 and B16 cells. Cell movement and migration to the denuded area was measured after 24 hours. Photos were taken at specific time points, as shown in Figure 10. Tumor cells showed differences in migration and invasion abilities after treatment with the four different drug formulations. The scratched gap had almost disappeared in control group by 24 hours’ incubation. Free Cur slightly inhibited the migration of cells compared with the control group, with migration rates of 45.68% and 58.9% in BEL7402 and B16, respectively. Free CA4P also played a minimal role in inhibiting tumor-cell migration. However, Cur + CA4P showed improved migration inhibition, with rates of 38.29% and 47.28% in BEL7402 and B16, respectively (Figure 10C and D). This indicated that combination treatment of Cur and CA4P can increase migration-inhibitory ability over single-Cur therapy. The combination-treatment therapy was superior to the single drug. Obviously, Cur-CA4P LPs and Cur-CA4P/GA LPs showed stronger inhibition efficacy on cell migration than the free mixture of Cur and CA4P. The possible cause was that LPs, as a carrier, increased cell internalization of drugs. Compared with Cur-CA4P LPs,
cell migration was remarkably inhibited for Cur-CA4P/GA LPs, with migration rates of only 15.74% and 15.69%, respectively. The results of the cell-migration test were consistent with cytotoxicity test results, where the Cur-CA4P/GA LPs group showed highest cell killing ability ($P<0.01$). GA molecules increased the active targeting of LPs and increased the cellular uptake of drugs. Therefore, Cur-CA4P/GA LPs inhibited the migration of cancer cells significantly.
In vivo real-time imaging of GA LPs
To investigate liver-targeting properties of GA LPs, we injected free DiR and DiR-GA LPs into H22 tumor-bearing mice. Results showed that NIRF signals of the free-DiR group were lower than the DiR-GA LPs group from 1 to 48 hours (Figure 11). No fluorescence signals in the tumor regions were detected for the free-DiR group, except at 2 hours. By contrast, DiR-GA LPs increased in accumulation in tumor regions and signals were sustained at 48 hours after injection. The GA LP drug loading system increased the accumulation of DiR in tumors. We concluded that the retention time of LPs were significantly prolonged in blood circulation. Moreover, GA improved the targeting of LPs and increased the accumulation of DiR in tumors. Therefore, the preferable active targeting ability of LPs to tumor sites will provide a key prerequisite for the superior combination therapy of Cur and CA4P.

In vivo therapeutic efficacy
In vivo antitumor efficacy of Cur-CA4P/GA LPs for H22 tumor-bearing mice was tested for 2 weeks. As demonstrated in Figure 12A, body weight in the free-Cur group showed an evident decreasing trend compared with the control group, indicating the systemic toxicity of free Cur. Interestingly, no evident weight difference between LP groups and the control was observed, suggesting that LP formulations can reduce systemic toxicity. In Figure 12B, mice treated with physiological saline displayed a rapid increase in tumor size during the experimental period, and those treated with the four drug formulations showed a slight increase, indicating that these drug formulations inhibited the development of tumors. Moreover, compared with single-drug treatment, the combination of Cur and CA4P was more effective in antitumor therapy, where the two drugs achieved an additive effect by proapoptotic and antiangiogenic activities.\(^\text{30}\)
Compared with free Cur + CA4P, the two LP formulations showed stronger antitumor efficacy (Figure 12B–D), which was consistent with MTT assays. The result was due to the fact that LPs improved the accumulation of Cur and CA4P in tumor regions via the EPR effect, resulting in high inhibition efficiency. Furthermore, as shown in Figure 12D, the tumor-inhibition rate of Cur-CA4P/GA LPs was 90.5%, which was higher than the Cur-CA4P LPs. A possible explanation was that GA LPs were taken up by tumor cells via GA-receptor-mediated endocytosis, which increased the antitumor effect by active liver-targeted delivery.

**Histochemical staining**

H&E staining was conducted to evaluate the antitumor effect of Cur-CA4P/GA LPs further. As shown in Figure 13, cancer cells showed no significant necrosis in the saline control group, in which nuclei were maintained in their full state. By contrast, tumor cells treated with drug formulations exhibited evident morphologic changes, with shrunken nuclei and low cellular density, suggesting inhibition in the development of tumor cells. Compared with the free-Cur group, the combination group produced more nuclear shrinkage, suggesting that the combination of Cur and CA4P achieved enhanced antitumor effects by proapoptotic and antiangiogenic activities. Moreover, tumor cells treated with Cur-CA4PLP and Cur-CA4P GALP formulations showed lower cellular density and more nuclear shrinkage in comparison with the Cur + CA4P group, indicating more drugs accumulated in tumor regions under treatment with the LP formulation. Among all the formulations, Cur-CA4P/GA LPs showed the most potent efficacy. This was probably due to the fact that GA LPs effectively delivered drugs to the tumor microenvironment via the EPR effect and increase drug uptake by tumor cells via GA-receptor-mediated endocytosis.

**Figure 12** (A) Mice were weighed at the same time. (B) Mice were killed and tumor blocks removed for photography. (C) Tumor volume changes in H22-bearing BALB/c mice after intravenous injection with normal saline and various drug-preparation groups. (D) Tumor growth–inhibition rate was calculated according to the formula. Tumor growth was monitored and plotted as tumor volume. Tumor growth–inhibition rate was calculated by weighing the tumor mass. Cur-CA4P LPs vs control group, Cur-CA4P/GA LPs vs control group. *P<0.05, **P<0.01.

**Abbreviations:** Cur, curcumin; CA4P, combretastatin A4 phosphate; LPs, liposomes; GA, glycyrrhetinic acid.

**Figure 13** Histological analyses of tumor tissue using H&E staining.

**Notes:** Scale bar 100 μm. Red arrows highlight immunohistochemical characteristics.

**Abbreviations:** LPs, liposomes; GA, glycyrrhetinic acid; CA4P, combretastatin A4 phosphate; Cur, curcumin.
Conclusion
In this work, novel Cur-CA4P/GA LPs were developed for effective antihepatoma therapy. GA LPs promoted cellular uptake and achieved more accumulation in tumor regions compared with GA-free LPs. Moreover, Cur-CA4P/GA LPs exhibited enhanced in vitro cytotoxicity and in vivo antitumor effects. In summary, GA-modified LPs for codelivery of proapoptotic and angiogenic drugs is a promising strategy for cancer therapy.

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Disclosure
The authors report no conflicts of interest in this work.

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